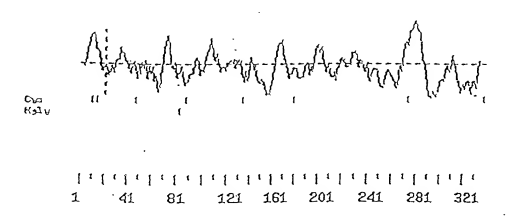
GG	AGTO	GACC	cxcc		CCC A	GGGC7	rgage	AACC	H OTA	S TCT	, CCY 5	rec	ccc	T ACC	GCC	CIC	F	·C TGT	L CTT	11 68
	CT			G G G G	R COI	.V.	q CCA	λ ∵GCG	CYC O	S AGT	G GGA	P CCG	L CTC	p P	K AAG	. CCC	s rcç	r Crc	CYC O	31 128
	L r CT		S. DAC				.p						ጥ አርር	L CTĆ	R CGG	C TGC	Ω CAG	G GGA	. P CÇT	51 168
	G G GG	V GTG	_		Y TAC		L CTG	· E GAG		L CTG		S TCC	S AGC	'R AGG	Y TAC		D GAT	Q CAG	, ACA	71 246
V GT		F TTC	_	P CCC		H ATG	K AAG	yèy K				G GGA	P. CGC	Y TAC				Y TAC		91 308
YYC N		S AGO	_		\$ 100	CIC P	CCC B	S S			L CTG	E GAG	L CTC		A GCC		G GGA	·V ·		111 368
SCC.		. C CC		L CTC	S TCA	. GCC	Q CAG	DCCC	GGC	P CCG	A GCG	V GTG	s rcc	S TCA	G GGA	G GGG	D. GYC	V GTA	т ЭСС	131 428
L CTA	CAG	C TGT	CYC G	T CT		Y TAT	G GGC	F TTT	D GAC	CAA	F TTT	A GCT	L CTG	Y	K XXG	E GAA	G GGG	D GAC	P CCT	151 <88
Х GCG	C CC	Y TAC	K AAG	H TAK	P CCC	E GAG	R AGA		Y TAC	R CGG	A GCT	S Agt	F TTC	CCC	I ATC	I ATC	Y VCG	V GTG	Т ЭСС	171 548
A GCC	A GCC	EAC TH	s Agc	G GGA	T ACC	Y TAC	R CGA	C TGC	Y TAC	S AGC		, s TCC	S AGC	R AGG	D GAC	P CCA	Y TAC	L CTG	W TGG	191 608
200 200	A GCC	P CCC	s Agc	QYC D	CCC P	L CTG	E GAG	L CTT	V GTG	V GTC	T ACA	G GGA	T ACC	S TCT	V GTG	YCC 2004	P CCC	S AGC	R CGG	211 668
L TTA	P ACOA	T ACA	E GAA	P CCA	PCCT	s TCC	S TCG	V GTA		E GAA	iic E	S TCA	E GAA	A GCC	T ACC	A GCT	E GAA	L CTG	ACĆ	231 728
V GTC	s TCA	F TTC	YCY	yyc N	XX.	V GTC	F TTC		.T ACT		T ACT	S TCT	'R Agg	s. Agt	I ATC	T DOA	T ACC	s Agt	CCY .	788
	E Gag	s TCA		S TCT	CCY B	A GCT	G GGT	P CCT			CYC CYC	Y TAC	TAC		X ANG	е с ес	n Aac	CLC	orc	271 8 48 ·
R CGG	1 . XTX	C TGC	L CTC	G GGG	A GCT	V GTG	I ATC	L CTA	I Ata	YIC I	L CIG	A GCG	G GGG	F TTT	CIG C		G/Q		i H TGG	291 908
CYC H	s NGC	R CGG	r Agg	X XXG	R CGC	L CTG	.R CGG	CYC	R Agg	G GGC	r Agg	A GCT	V GTG	ĆYG Ó	r Agg	P CCG	L CTT	CCG	CCC B	311 968
	CCCC E	CCC	CTC L	P CCC	cyc G	T ≫C	R CGG	X XXX	S TCA	cyc H	G GGG	G GGT	CYC C	d Gat	G GGA	G GGC	·R CGA	CAG Q	D GAT	331 1028
	EYC SYS	e Ngc	R CGC	G GGG	L TTA	C TGT	s TCA	TGA					•	-						340 1055

FIGURE 1a

CCGCTGAACCCCAGGCACGGTCGTATCCAAGGGAGGGATCATGGCATGGGAGGCGACTCAAAAGACTGGCGTGTGTGGAG	1134
COTOGRAGOLOGICAGAGGCTACAGCTGTGGAAACGAGGCCATGCTGCCTCCTCCTGGTGTTCCATCAGGGAGCCG	1213
TTCGGCCAGTGTCTGTCTGTCTGCCTCTCTGTCTGAGGGCACCCTCCATTTGGGATGGAAGGAA	1292
CCCATCCTCCTCCCTGCACACTGTGGATGACATGGTACCCTGGCTGG	1371
ANTATGGGCTCCAGACGGATCTCTAAGGTTCCCAGCTCTCAGGGTTGACTCTGTTCCATCCTCTGTGCAAAATCCTCCT	1450
GTGCTTCCCTTTGGCCCTCTGTGCTCTTGTCTGGTTTTCCCCAGAAACTCTCACCCTCACTCCATCTCCCACTGCGGTC	1529
TAACAAATCTCCTTTCGTCTCTCAGAACGGGTCTTGCAGGCAG	. 1608
AGCACGTTGCCCGCTTCCCTTCACATTAGAAAACAAGATCAGCCTGTGCAACATGGTGAAACCCTCATCTCTACCAACAA	1687
MCAAAAAAACACAAAAATTAGCCAGGTGTGGTGGTGCATCCCTATACTCCCAGCAACTCGGGGGGCTGAGGTGGGAGA	1766
ATGGCTTGAGCCTGGGAGGCAGAGGTTGCAGTGAGCTGAGATCACACCACTGCACTCTAGCTCGGGTGACGAAGCCTGA	1845
CCTTGTCTCAAAAAATACAGGGATGAATATGTCAATTACCCTGATTTGATCATAGCACGTTGTATACATGTACTGCAAT	1924
ATTGCTGTCCACCCCATAAATATGTACAATTATGTATACATTTTTAAAATCATAAAAATAAGATAATGAAAAAAAA	2003
WWWWWWGGGCGGCCGCTAGACTAGTCTAGAGAACA	2047

FIGURE 1b



MSPSPTALFCLGLCLGRVPAQSGPLPKPSLQALPSSLVPLEKPVTLRCQGPPGVDLYRLE KLSSRYQDQAVLFIPAMKRSLAGRYRCSYQNGSLWSLPSDQLELVATGVFAKPSLSAQP GPAVSSGGDVTLQCQTRYGFDQFALYKEGDPAPYKNPERWYRASFPIITVTAAHSGTYRC YSFSRDPYLWSAPSDPLELVVTGTSVTPSRLPTEPPSSVAEFSEATAELTVSFTNKVFT TETSRSITTSPKESDSPAGPARQYYTKGNLVRICLGAVILIILAGFLAEDWHSRRKRLRH RGRAVQRPLPPLPPLPOTRKSHGGQDGGRQDVHSRGLCS

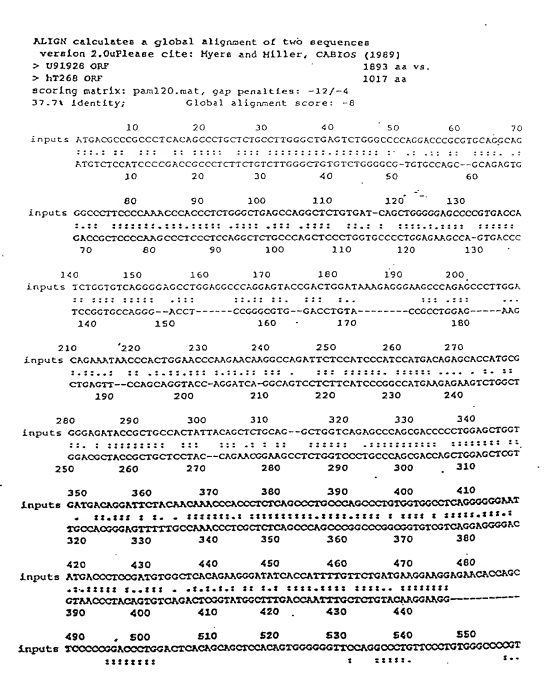


FIGURE 3a

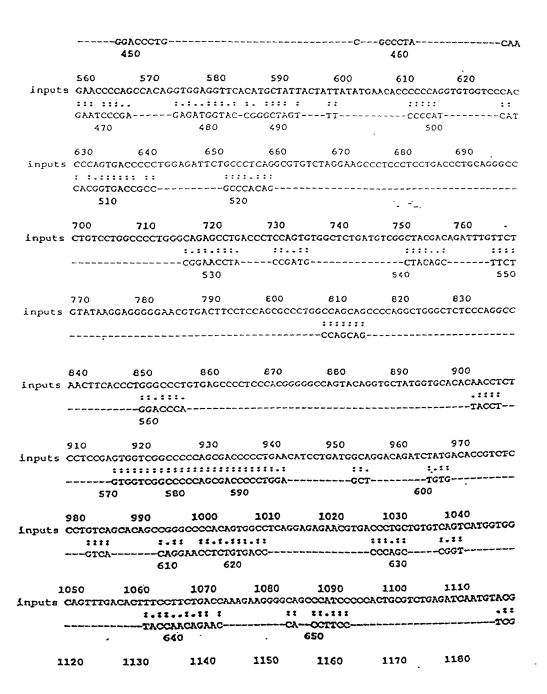


FIGURE 3b

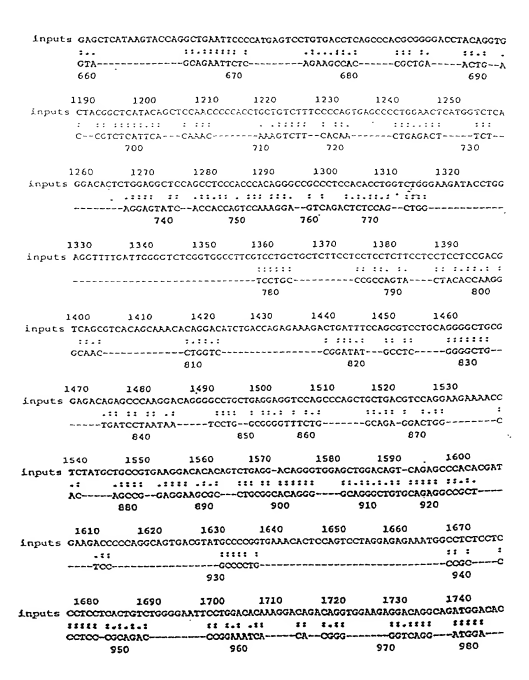


FIGURE 3c

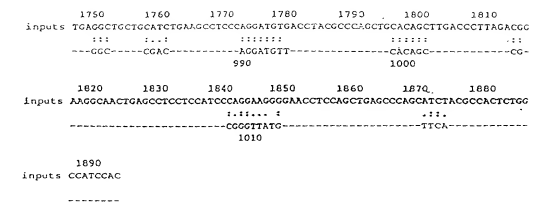


FIGURE 3d

```
ALIGN calculates a global alignment of two sequences
 version 2. OuPlease cite: Hyers and Hiller, CABIOS (1989)
 > GenPept U91928 - Human clone HL9 monocyte inhib
 scoring matrix: pam120.mat, gap penalties: -12/-4
 23.01 Identity; Global alignment score: -642
                20
                      30
                            40
                                  . 50
Inputs HSPSPTALFCLGLCLG-RVPAQSGPLPAPSLQALPSSLVFLERPVTLRCQGPPGVDLYRLEKLSSS----
     HTPALTALLCLGLSLGPRTRVQAGPFPKPTLWAEPGSVISWGSPVTIWCQGSLEAQEYRLDKEGSPEPLD
             20 30 40 50
                                 110 . . . 120
                          100
               80
Inputs RYQ----DQAVLFIPAHKRSLAGRYRCSYQNGSLHSLPSDQLELVATGVFAKPSLSAQPGPAVSSGGDV
    RUNPLEPKNKARFSIPSHTEHHAGRYRCHYYSSAGHSEPSDPLELVHTGFYNKPTLSALPSPVVASGGNH
             90 100 110 120 130. 140
TLRCGSQKGYHHFVLMKEGEHQLPRTLDSQQLHSGGFQALFPVGPVNPSHRWRFTCYYYYMNTPQVWSHP
        150 160 170 180 190 200 210
                           140
inputs -----GFDQFALYKEGDP-----
                            ............
    SDPLEILPSGVSRKPSLLTLQGFVLAPGQSLTLQGGSDVGYDRFVLYKEGERDFLQRPGQQPQAGLSQAN
        220 230 240 250 260
inputs ------ERW--
          . . . .
    ftlgpvspshggqyrcygahnlssehsapsdplnilmagqiydtvslsaqpgptvasgenvtllcqshhq
        290 300 310 320 330 .340
                                    190
                       170
                              180
inputs -----YRASFPIITVTAAHSGTYRCYSFSSRDPYLWSAPSDPLELVVTG
              fotflltkegnahpplrlrshygnhkygnefphspytsnhagtyrcygsyssnphllsfpseplelhysg
        360 370 380 390
                                400 410
                             240
            220
                 230
inputs tsytpsrlpteppss--vaefseataeltysftnkvf-----ttetsrsittspkesd--spagpa-
    ESGGSSLPPTGPPSTPGLGRYLEVLIGVSVAFVLLLFLLLFLLLRRQRHSKHRTSDQRKTDFQRPAGAAE
                                     460
             440 450 460 470
        430
             280
                       290
       270
. . .::.
    . .::: .:.
```

FIGURE 4a

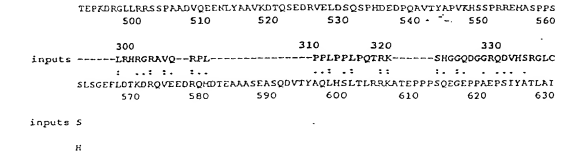


FIGURE 4b

```
Alignments of top-scoring domains:

ig: domain 1 of 2. from 41 to 90: score 4.1, E = 6.1

->GesvtLtCsvsgfgppgvsvtWyfkngk.lgpsllgysysrlesgek

hT268

41 EKPVTLRCQGP-----PGVDLY-RLEKISSS------RYQDQ--70

anlsegrfsissltLtissvekeDsGtYtCvv<-+

+tL i ++++G Y+C

hT268

71 --------AVLFIPAMKRSLAGRYRCSY 90
```

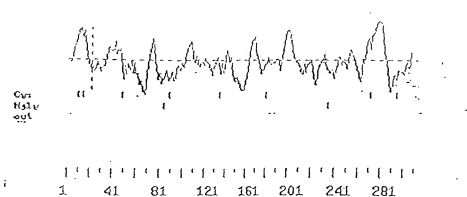
```
ig: domain 2 of 2, from 127 to 182: score 19.1, E = 0.1

*->GesvtLtCsvsgfgppgvsvtWyfkngk.lgpsllgysysrlesgek
G++vtL+C+++ + ++ y k+g++ + y+++
hT268 127 GGDVTLQCQTR---YGFDQFALY-KEGDpAP----YKNPERWYR-- 162

anlsegrfsissltLtissvekeDsGtYtCvv<--
++++i+v++ sGtY+C
hT268 163 -------ASFPIITVTAAHSGTYRCYS 182
```

GYC	TCGA	'cccı	vocco	TCCC	CTTC	ccre	CTTG	GCC I	LTAD.	.GCTC	:XGGX	CTG0	cric	CAGA	лсс	H ATG	S TCT	P CCA :	λ GCC	د 74
S TCA	CCC	т : ДС1	F 7. 770	-			· · · · · · · · · · · · · · · · · · ·					. Q						S Agt	G G	2 〈 13 〈
	CLC T	. CCC	X SXA	P CCT	S TCC	L	Q CAG		CYC G		5 3.57	s ncc	t CTG	V GTA	CCC L	L CTG	G GGT	CY2 G	S TOA	· 4 4
V	I ATT	L CTG	A DOK	C	_	G GGA	CCT P			v GTG		L TTA	У ТАТ	R CGC	L CTG	E GAG	X AAA	L CTG	K KK	6 ¢ 25 ¢
CCG		X Dag	Y TAT				D GAC					CCY 5	YCC T	YIC H	GYY E	R AGA		14.	ъ GCТ	8 (31 4
	R CGG	Y TAT	R CGA	JCC C		Y Tat	-	11 T.K.S.			CYC H	H TGG	S TCT		P CCA		. Þ GAC	Q CAG	L CTT	10¢ 37¢
E GAG	L CTA	I TTA	Х GCT	T AOA	G GGT	V 670	Υ ΤλΤ		K KK			L CTC	S TCA	k GCT	H CAT	P CCC	s S	S AOT	A GCA	124 434
V GTC	P CCT	CYY CYY	G GGC	R AGG	D GAT		T ACT	-	K AAG		_	S AGC	P CCA	Y TÀC	s አርፕ	F TIT	D GAT	GYY E	F TTC	1 < < < 9 <
v GTT		Y TAC	K XXX	E GAA	G GGG		T ACT	_	P CCT	-	K N.S	R Aga	P CCT	E GAG	к х хх	ri TGG	Y TAC	R CGG	, ССС	164 554
n Aat	F TTC	P CCC) J	I ATC	T ACA		T ACT		K GCT		S Agt	-	T ACG	Y TAC			Y. TAC	•	F TTC	184 614
s TCC	s XGC	S TCA	Ś TCT	P CCA	Y TAC	L CTG	w TGG	S TCA	A GCC	cće Ł	S Agt		· p CCT	L CTA	V GTG	L CTT	ėle V	V GTT		204 674
G GGA	L CTC	S TCT	SCC Y	T ACT	CCC P	S AGC	CYC O	V GTA	CCC P	T ACG	E GAA	GYY E	S TCA	F TTT	P CCT		T ACA		s rcc	224 734
s rcc.	r XG G	R AGA	P CCT	S TCC	I ATC	L TTA	P CCC	T ACA	N AAC	K XXX	I ATA			T ACT	E GAA	K AAG	P	H ATG	И Тал	794
I ATC .	T ACT	ecc Y	S TCT	CCY D	E GAG	GGG	CLC F	s NGC	P CCT	ς CCλ	I Att		F TTT			CYC O		X. TAT		264 854
X XXG	G GGG	N Aat	L CTG	V. GTC	R CGG	I Ata	C AGC	L CTT	G GGT	ecc Y	YCC T	I TTA	I ATA	I ATA	I ATT	L TTG		GGG G·		284 914
L CTA	A GCA	E GAG	D GAT	H TGG	CAC	S Agt	R CGG	X Aag	, XX	TGC C	L CTG	cyr	CAC	r Agg	H ATG	r aga		. L .TTG		304 974
AGG (CCY	CTA		CCC	CTC	CCA														314
1083 GGACATACTCAAGAGGGAGGGATTGACCAGACATCCATGCACAACCATGGACATCACCACTAGAGCCACAGACAT 1083																				
A.			.5641	,			unn									-				1163

FIGURE 6



MSPASPTFFCIGLCVLQVIQTQSGPLPKPSLQAQPSSLVPLGQSVILRCQGPPDVDLYRL EKLKPEKYEDQDFLFIPTMERSNAGRYRCSYQNGSHWSLPSDQLELIATGVYAKPSLSAH PSSAVPQGRDVTLKCQSPYSFDEFVLYKEGDTGPYKRPEKWYRANFPIITVTAAHSGTYR CYSFSSSSPYLMSAPSDPLVLVVTGLSATPSQVPTEESFPVTESSRRPSILPTNKISTTE KPMNITASPEGLSPPIGFAHQHYAKGNLVRICLGATIIILLGLLAEDWHSRKKCLQHRM RALQRPLPPLPLA

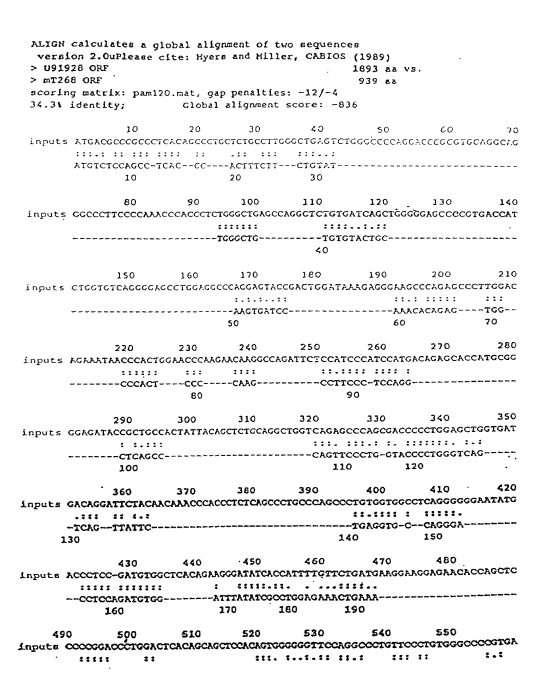


FIGURE 8a

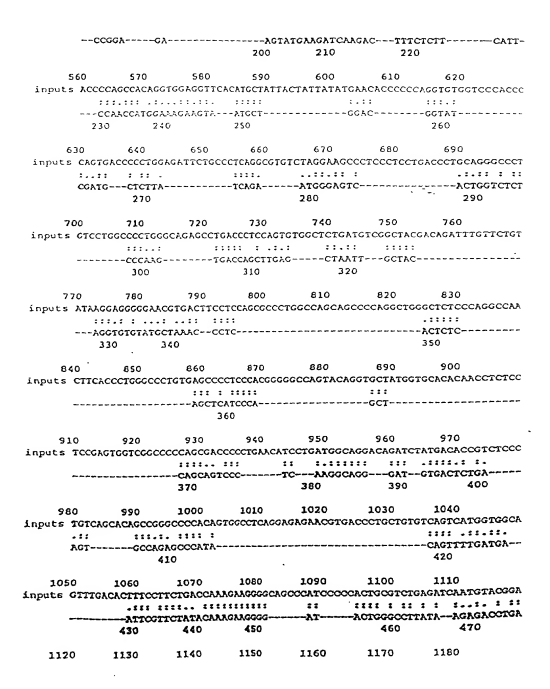


FIGURE 8b

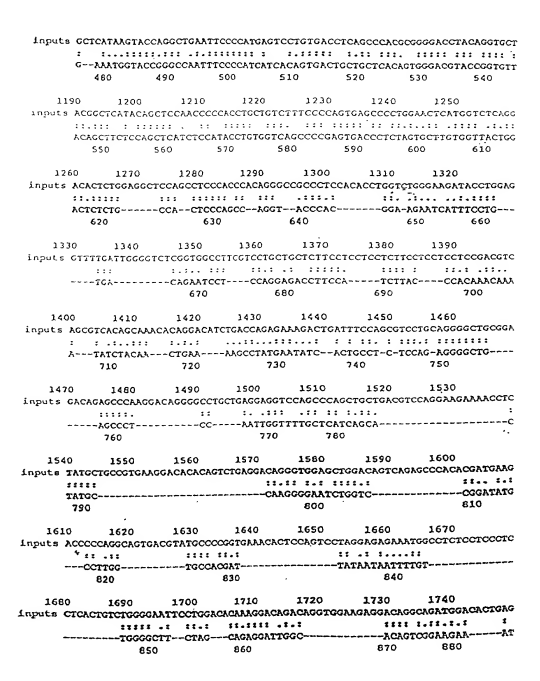


FIGURE 8c

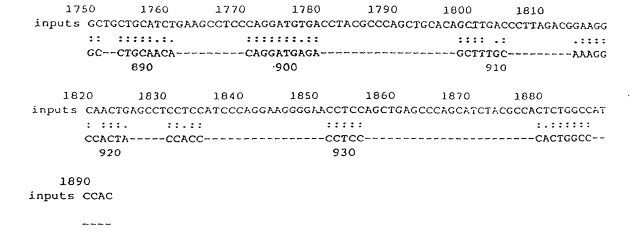


FIGURE 8d

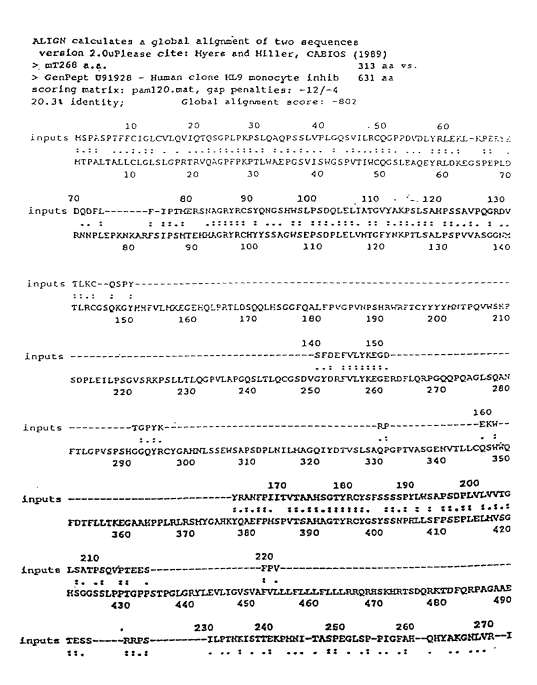


FIGURE 9a

	TEPKDRGLLRE	RSSPAADVQEEN	LYAAVKDTQSI	EDRVELDSQSP	YTVAQ9D3@H	APVEHSSPRI	REMASPPS
	500	510	520	530	540	550	560
						~	
	280	290		3	00		310
inputs	CLGATIIIILL	.GLLAEDWH		srkkclqh	RHRALQRPL-	PP	LPL
	:	::		:		::	:
	SESCEFLOTKD	RQVEEDRQHDT	PARSEASQOV	/TYAQLHSLTLI	rrkateppps(egeppaeps	IALTAYI
	570	580	590	600	610	620	630
inputs	7.						
	H						

FIGURE 9b

ig: domain 2 of 2, from 128 to 183: score 9.6, E = 1.6

->GesvtLtCsvsgfgppgvsvtWyfkngk.lgpsllgysysrlesgek
G +vtL C++ ++ y k+g++ + y+r+e +
GRDVTLKCQSP---YSFDEFVLY-KEGDtGP-------YKRPEKW-Y 162

anlsegrfsissltLtissvekeDsGtYtCvv<-+
+ ++i++v++ sGtY+C

mT268 163 RA-------NFPIITVTAAHSGTYRCYS 183

```
· ALIGN calculates a global alignment of two sequences
 version 2.0uPlease cite: Myers and Hiller, CABIOS (1989)
 > hT268 a.a.
                                        339 Ac vs.
 > mT268 a.a.
                                        313 ca
 scoring matrix: pan120.mat, gap penalties: -12/-4%
'64.4% identity; Global alignment score: 1011
          . 10
                    20
                                          50.
                          30 - 40
                                                  60
 inputs HSPSPTALFCLGLCLGRV-PADSGPLPXPSLQALPSSLVPLEKPVTLRCQGPPGVDLYRLEKLSSSRYQD
      HSPASPTFFCIGLCVLQVIQTDSGPLPKPSLQAQPSSLVPLGQSVILRCQGPPDVDLYRLEKLKPEKYED
                                  40
                                          50
                                                 60
            10
                   20
                           30
            03
                    90
                          100
                                  110
                                          120
                                                 130
inputs QAVLFIPAYKESLAGRYRCSYQNGSLWSLPSDQLELVATGVFAKPSLSAQPGPAVSSGGDVTLQCQTRYG
      QDFLFIPTHERSNAGRYRCSYQNGSHWSLPSDQLELIATGVYAXPSLSAHPSSAVPQGRDVTLKCQSPYS
                                 110
                                         120
                                               130 -
            03
                   90
                         100
                          170
                                  180
                                          190
                                                 200
            150
                   160
inputs FDQFALYKEGDPAPYKNPERWYRASFPIITVTAAHSGTYRCYSFSSRDPYLWSAPSDPLELVVTGTSVTP
      FDEFVLYKEGDTGPYKRPEKWYRAI#PIITVTAAHSGTYRCYSFSSSSPYLWSAPSDPLVLVVTGLSATP
                                 180
                                         190
                          170
                  160
                  230 240
                                  250
                                          260
           220
inputs splatepassvaefseataeltysfthkyfttetsrsittsakesdsaagarqyytkgnlyriclgavi
      SQVPTEESFPYTESSRRPSILP---TNKISTTEKPMNITASPEGLSPPIGFAHQHYAKGNLVRICLGATI
                                    250
                            240
                  230
    280
                                  320
                                          330
                           310
            290
                   300
inputs LIILAGFLAEDWHSRRKRLRHRGRAVQRPLPPLPPLPQTRKSHGGQDGGRQDVHSRGLCS
      IIILLGLLAEDWHSRKKGLQHRWRALQRPLPPLP-LA------
                     300
                          310
             290
```

FIGURE 11

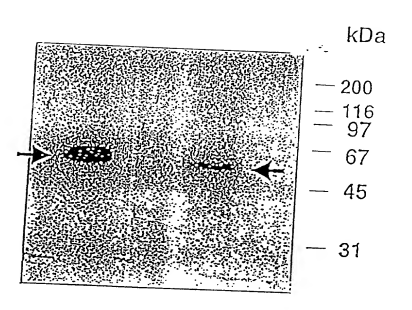


FIGURE 12

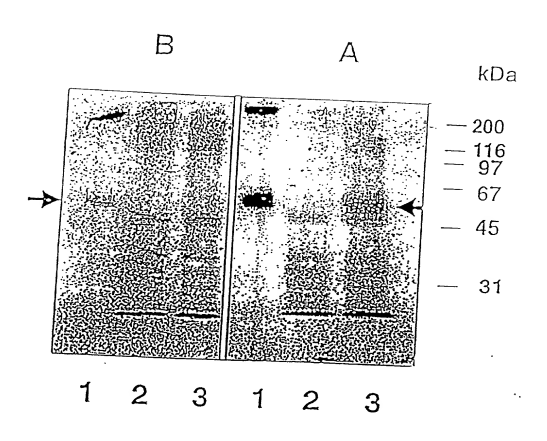


FIGURE 13

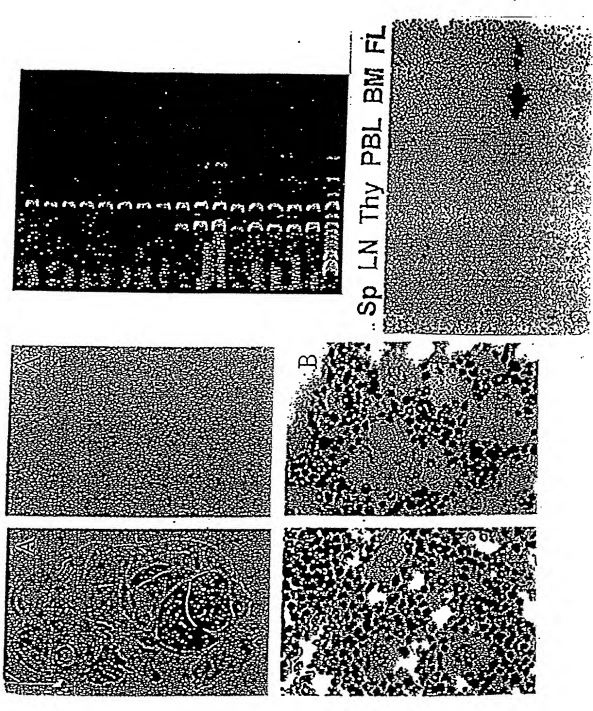
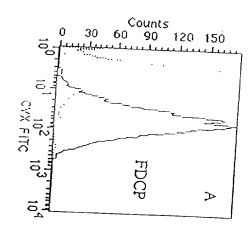
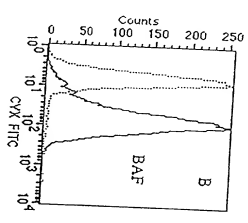


FIGURE 14





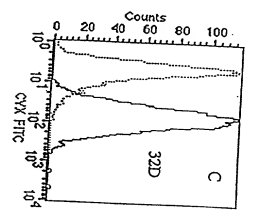


FIGURE 15

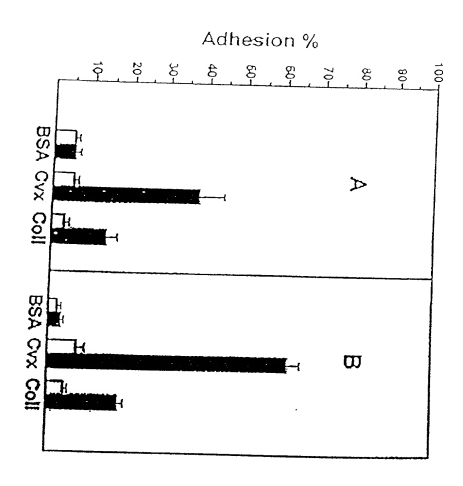


FIGURE 16 ·

1

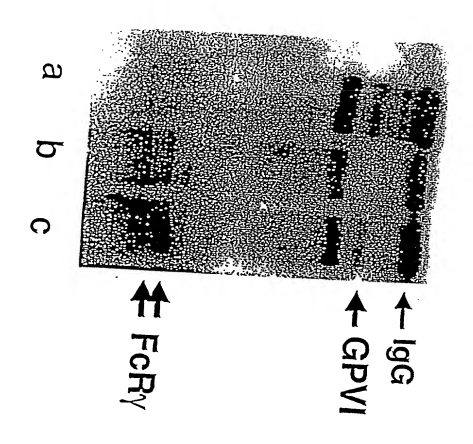


FIGURE 17

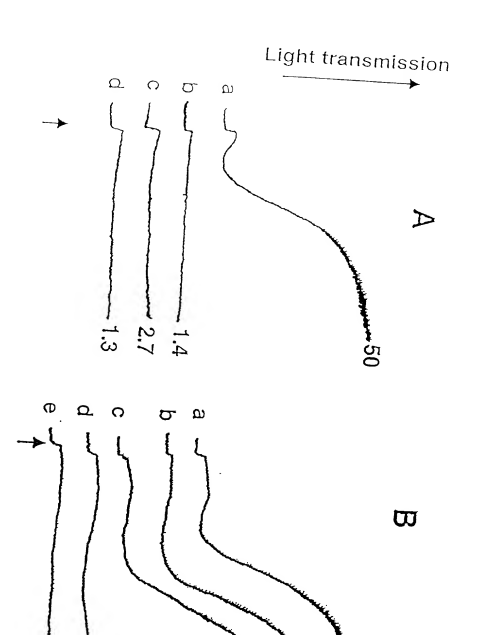
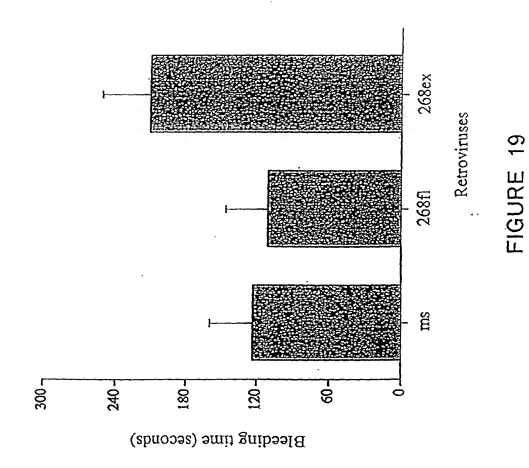


FIGURE 18

<u>ශ</u>

<u></u>





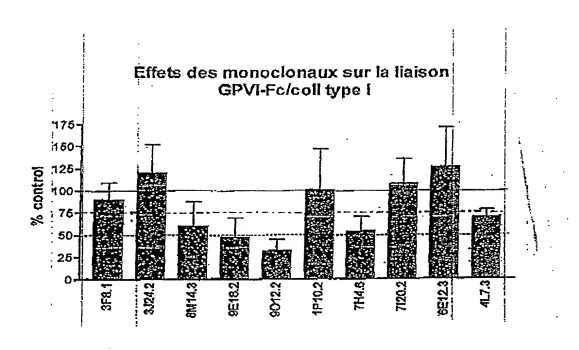
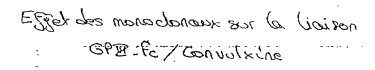


FIGURE 20



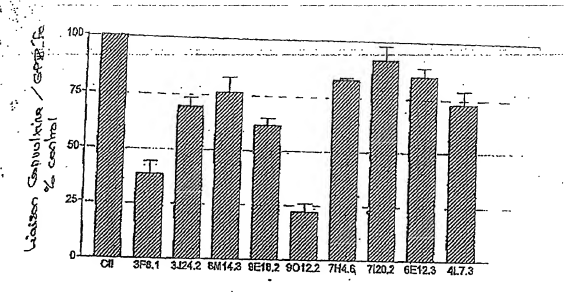
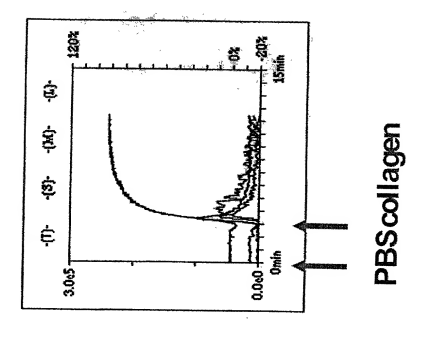


FIGURE 21

(Sheet 33 of 42)



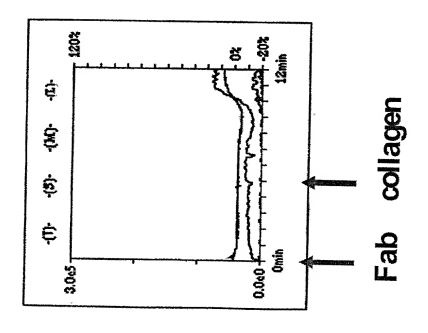


FIGURE 22

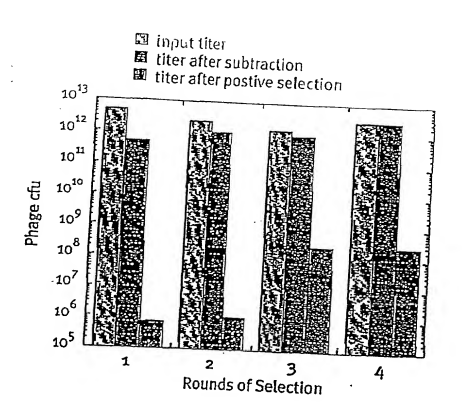


FIGURE 23

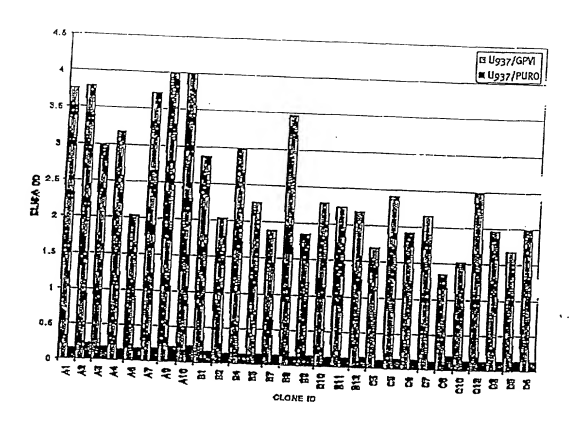


FIGURE 24a

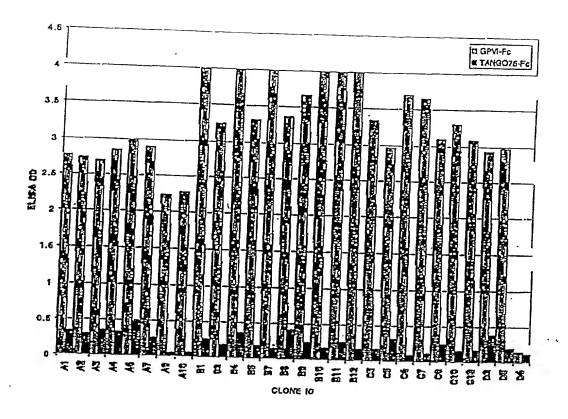


FIGURE 24b

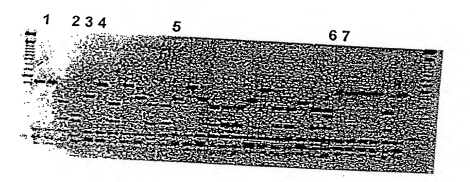


FIGURE 25

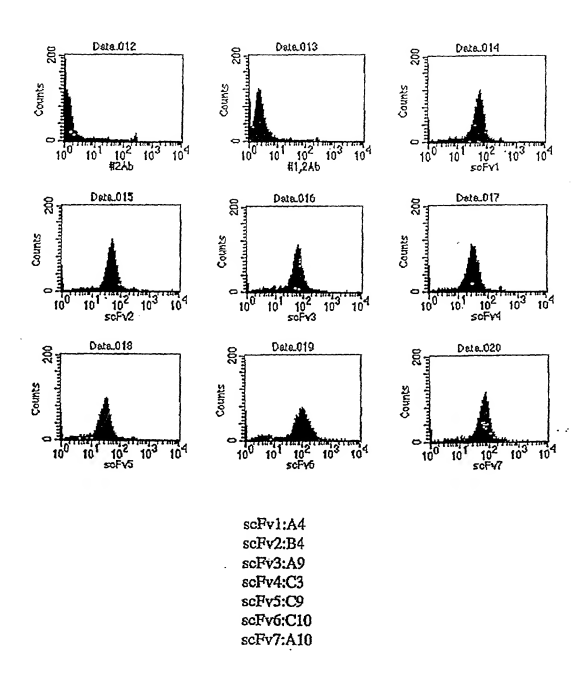


FIGURE 26

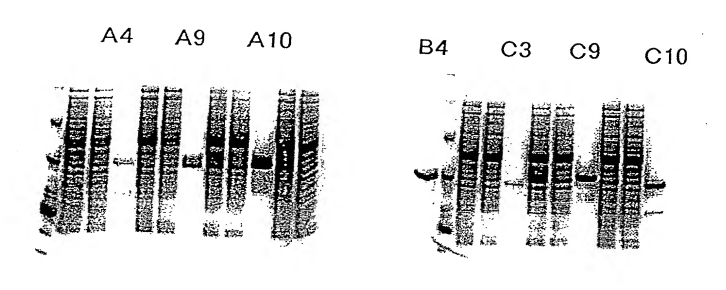


FIGURE 27

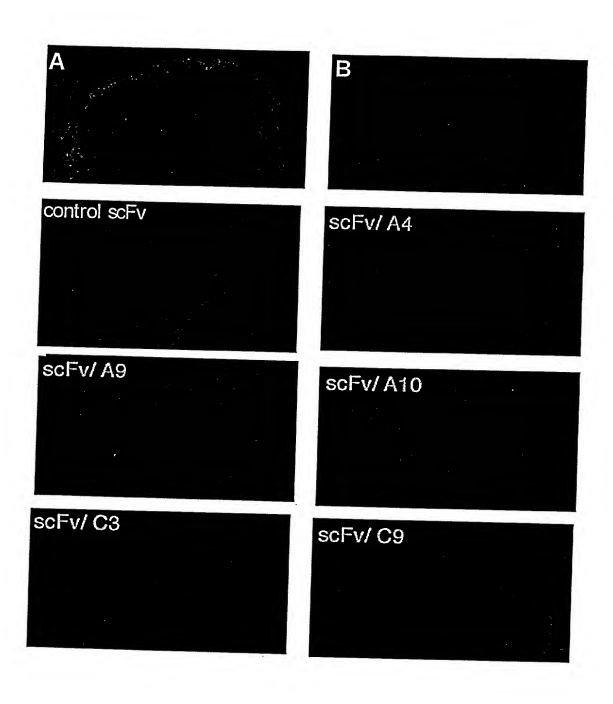


FIGURE 28

